

VistaSeq® Breast Cancer Panel

Specimen ID:

Control ID: Acct#: Phone:

**TESTING** 

Patient Details

DOB: Age (yyy/mm/dd):

Gender: Patient ID: Specimen Details

Date collected: Date received:

Date entered: Date reported: **Physycian Details** 

Ordering: Referring: ID:

NPI:

# **POSITIVE**

At least one clinically significant variant was detected.

## **RESULTS AND INTERPRETATION**

	GENE	CLASSIFICATION	ZYGOSITY	VARIANT DETECTED	AMINO ACID CHANGE	CANCER RISK
+	ATM	LIKELY PATHOGENIC	Het	c.237delA	p.Lys79fsX37	MODERATE

**Variant Summary:** A heterozygous c.237delA (p.Lys79fsX37) likely pathogenic variant was detected in exon 4 of ATM. This frameshift variant is predicted to result in a downstream premature termination codon and has been previously reported in ClinVar and in the literature. Therefore, this variant has been classified as likely to be associated with an increased risk for breast and pancreatic cancer. (NM\_000051; hg19 chr11:g.108099955-108099956)

ATM (ATM serine/threonine kinase; OMIM 607585) is a protein kinase that has roles in normal cell cycle control, development of the nervous and immune systems, and coordination of DNA damage responses promoting genomic stability. Heterozygous germline ATM mutations have been identified and associated with familial breast and pancreatic cancers. Biallelic germline ATM mutations can cause ataxia-telangiectasia, a progressive neurological/immunodeficiency disorder with a predisposition to cancer development, including leukemias and lymphomas.

### **Clinical Significance: Moderate Cancer Risk**

This mutation is clinically significant and is associated with an increased cancer risk. Current NCCN guidelines for ATM carriers suggest annual mammography and consideration of breast MRI with contrast starting at age 40. Risk reduction management should be based on family history (www.nccn.org). In addition to this individual being at increased risk, other family members may also be at risk. There is a 50% (1 in 2) chance of a first-degree relative having this mutation. Please call (800) 345-4363 to speak to a Labcorp Genetic Counselor to discuss if targeted analysis for other family members is appropriate.

# This result is associated with the following cancer risks:

Lifetime Increased Risk 24-48% Female breast

Lifetime Possible Increased Risk Pancreatic

\*See table below for additional risk information

## **RECOMMENDATIONS**

Genetic counseling is recommended to discuss the clinical implications of this result. Genetic counselors are available for health care providers to discuss this result further at (800) 345-GENE. To refer your patient for genetic counseling through Integrated Genetics, please call the scheduling line at (855) 422-2557.



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Patient: TESTING DOB:		Patient ID:			Control ID:			Specimen ID Date collected				
CANCI	ER TYPE	CANCER RISK			RISK FOR GENERAL POPULATION		RELATED TO					
Br	east											
To a	To age 70		24-48%		12% Females		ATM					
Pan	Pancreatic											
To age 70		Risk not yet determined			1.5%		ATM					
LIST OF ALL GENES IN PANEL												
ATM	CDH1	BARD1	BRIP1		RAD50	MRE	11A	RAD51D				
NBN	PTEN	BRCA1	CHEK2		STK11	RAD	51C	FAM175A				
NF1	TP53	BRCA2	PALB2		MUTYH (biallelic)							

# **ADDITIONAL INFORMATION**

Specimen Type: Whole Blood

Indication for Testing: The indication for testing for this patient is a reported personal and/or family history of breast and pancreatic cancers.

Variant Classification: Variant classification is a weighted assessment that incorporates but is not limited to the following components: prevalence of a variant in the unaffected (general) population, evidence of co-segregation in affected individuals, review of locus specific databases and observed/reported co-occurrence with other deleterious variants within the gene, published functional evidence linking a variant to phenotypes, and predicted functional impact as determined using in-silico analyses. Variants classified within each gene are reported in accordance to the ACMG standards and guidelines. Evidence affecting a variant classification that alters its clinical significance will be reported via an amended report. Pathogenic variants negatively affect normal gene function, are associated with disease, and should be used in clinical decision making. Likely pathogenic variants are strongly suggestive of normal gene function being negatively affected, and when combined with other evidence of cancer, may be used in clinical decision making. Variants of uncertain significance (VUS) have unknown effects on gene function, have not been previously reported or have been reported with inadequate or conflicting evidence regarding pathogenicity, clinical relevance, or cancer risk. A VUS should not be used in clinical decision making but additional monitoring may be considered. Likely benign variants are strongly suggestive of having no effect on gene function and are unlikely to have an increased risk for cancer. Benign variants have sufficient evidence to be considered of no clinical significance. Likely benign, benign and synonymous variants are not reported, but are available upon request.

#### METHODOLOGY AND LIMITATIONS

Next generation sequencing is used to examine the entire gene coding regions, as well as flanking non-coding regions, of genes known to be involved in the development, progression, and susceptibility of cancer. Flanking regions for the BRCA1 and BRCA2 genes include +/- 20bp and +/-10bp for all other genes. Copy number variations are assessed by microarray or multiple-ligation-probe amplification assay (MLPA) to detect gross deletions and duplications. Due to inherent limitations in the sequence analysis methods used, some variants may be missed. The presence of pseudogenes can interfere with the ability to detect variants in certain genes. Results are reported using nomenclature recommended by the Human Genome Variation Society (HGVS http://www.hgvs.org/). Each gene sequence is interpreted independently of all other gene sequences. However, variants in different genes may sometimes interact to cause or modify a typically monogenic disease phenotype. The occurrence of cancer due to genes not analyzed with this test is possible. Additional details regarding technical specifications and limitations of this assay are available on our websites, www.labcorp.com, www.integratedgenetics.com, and www.integratedoncology.com.

This test was developed and its performance characteristics determined by LabCorp. It has not been cleared or approved by the Food and Drug Administration.

Date Issued: FINAL REPORT Page 2 of 3





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## **REFERENCES**

- 1. National Comprehensive Cancer Network. Clinical practice guidelines in oncology, genetic/familial high-risk assessment: breast and ovarian. Available at: www.nccn.org. 2010. Accessed 5.29.13.
- 2. Rehm H. et al. Working Group of the American College of Medical Genetics and Genomics Laboratory Quality Assurance Commitee. ACMG clinical laboratory standards for next-generation sequencing. Genet Med. 2013 Sep;15(9):733-47.
- 3. Tung N. et al. Frequency of mutations in individuals with breast cancer referred for BRCA1 and BRCA2 testing using next-generation sequencing with a 25-gene panel. Cancer. 2015 Jan 121(1):25-33.
- 4. LaDuca H. et al. Utilization of multigene panels in hereditary cancer predisposition testing. Genet Med. 2014 Nov;16(11):830-7.

# **Released By:**

#### PERFORMING LABORATORIES

TG LabCorp RTP 1912 T.W. Alexander Drive, RTP, NC 27709-0150 Lab: (800) 345-4363 Dir: Arundhati Chatterjee, MD For inquiries, the physician may contact the lab using the numbers indicated above.